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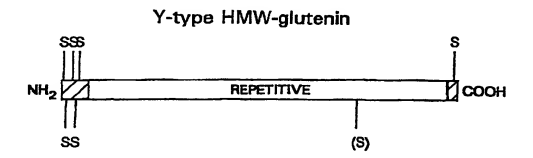
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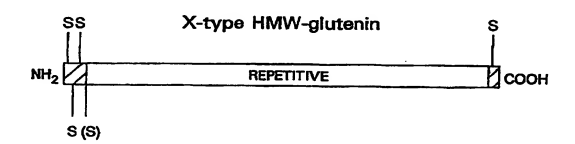
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(54) Title: GLUTENIN GENES AND THEIR USES





(57) Abstract

The present invention is directed to methods of improving flour by altering glutenin content in seeds of plants, particularly wheat. The invention relates to methods of introducing a recombinant construct comprising a glutenin gene into a parental plant.

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GLUTENIN GENES AND THEIR USES

Field of the Invention

The present invention relat's generally to plant molecular biology. In particular, it relates to nucleic acids and methods for improving glutenin content of plants.

BACKGROUND OF THE INVENTION

The glutenins, which include both high molecular weight (HMW) glutenin subunits and low molecular weight (LMW) glutenin subunits, comprise an economically important class of wheat seed storage proteins. The apparent molecular weights of the individual HMW glutenin polypeptides or subunits range from 90 to 200 kDa. These subunits crosslink by disulfide bonds among themselves and with LMW glutenin polypeptides to form polymers exceeding one million daltons in molecular weight. HMW glutenins constitute 8-10%, while LMW glutenins constitute 15-20% of the total endosperm protein. Both HMW and LMW glutenin proteins play important functional roles in determining the end-uses of wheat flour.

In wheat, HMW glutenins are encoded at the Glu-1 loci on the long arms of the group 1 chromosomes. Each locus consists of two separate genes, encoding an x-type and a y-type subunit, respectively. These pairs have never been confirmed to be separated by recombination. This has made determination of their separate contributions to bread dough difficult to assess by genetic correlation studies. For a the genetics and biochemistry of glutenin polypeptides, see, Shewry et al., J. Cereal Sci. 15:105-120 (1992).

Both the quantity and identity of specific HMW glutenin alleles contribute to the differences in bread-making quality of various cultivars. For instance, deletion of glutenin gen s results in a decrease in the overall levels of HMW glutenins, which results in decreases in bread-making quality (see, e.g., Lawrence et al. J. Cereal Sci 7:109-112 (1988)).

The effects f overproducing HMW glut nin on protein accumulation and baking quality has n t been assessed because such lines of what have not be n found among natural

populations. In addition, dir ct alteration of the glutenin subunits that form the polymers is not possible using standard breeding methods. Thus, the art lacks reproducible and fficient methods of producing lines with altered glutenin contents. The present invention addresses these and other needs.

SUMMARY OF THE INVENTION

The present invention provides methods of increasing glutenin in the endosperm of wheat plants. The methods and plants of the invention therefore are useful in providing flour and dough having improved end-use properties. The methods comprise introducing into a parental wheat plant a recombinant expression cassette comprising a nucleic acid encoding a glutenin polypeptide and selecting progeny wheat plant having increased glutenin content in the endosperm of mature seed. The glutenin content of the progeny is preferably at least about 15% greater than the glutenin of the parental wheat plant.

Any method of introducing the expression cassette into the parental plant can be used. Particle bombardment is a convenient method for producing transgenic plants. Once the expression cassette is stably integrated into the genome, standard sexual crosses can be used to introduce the expressi n cassette into desired lines.

In some embodiments, the nucleic acid introduced into the plant encodes a chimeric glutenin polypeptide. For instanc, the chimeric polypeptide may comprise sequences from an x-type glutenin polypeptide and a y-type glutenin polypeptide. Exemplary genes for this purpose include the Glu-Dl-lb gene and the Glu-Dl-2b gene.

The expression cassette may further comprise a sedspecific promoter to direct expression of the introduc d nucleic acid to the endosperm. A convenient promoter for this purpose is the promoter from the Glu-Dl-2b gene.

Any wheat cultivar can be used as the parental line in the present invention. An exemplary cultivar is Bobwhit .

The inventi n also provid s what plants comprising a recombinant expression cass the comprising a nucl ic acid encoding a glutenin polyp ptide. The plants of the invention

have a glutenin content in th endosperm of a matur s d at least about 15% greater than the glutenin content in the endosperm of a mature seed from a par ntal wheat plant. The percentage of total indosperm pr t in which is glutenin will usually depend upon the glutenin content of the parental line. Usually, HMW glutenins account for at least about 15% of th total protein in the endosperm of the mature seed from plants of the invention.

The invention further provides recombinant constructs comprising a wheat glutenin gene promoter of about 400 to about 2800 nucleotides operably linked to a heterologous polynucleotide sequence. These constructs are particularly useful in directing expression of the heterologous sequenc to seeds of transgenic plants. An exemplary wheat glutenin gene from which the promoters of the invention can be derived is Glu-D1-2b.

Definitions

The term "plant" includes whole plants, plant organs (e.g., leaves, stems, roots, flowers, etc.), seeds and plant cells and progeny of same. The class of plants which can be used in the methods of the invention is generally as broad as the class of higher plants amenable to transformation techniques, including both monocotyledonous and dicotyledonous plants.

A "heterologous sequence" is one that originates from a foreign species, or, if from the same species, is substantially modified from its original form. For example, a promoter operably linked to a heterologous structural gene is from a species different from that from which the structural gene was derived, or, if from the same species, one or both are substantially modified from their original form.

A "glutenin polypeptide" is a gene product of a glutenin gene or glutenin polynucleotide sequence. A glutenin polypeptide can be either a LMW glutenin or a HMW glutenin. A glutenin polypeptide contains cysteine residues by which disulfide bonds are formed with other glutenin polypeptides to form polymers. The composition and size f th repeat region is also important to polymer f rmation.

A "chimeric glutenin p lypeptide" is glutenin gene product that comprises a m dified amino acid sequence. Modifications, as explain d in detail below, can b in the form of substitutions, deletions, or additions of single amino acids or groups of amino acids. Thus, chimeric glutenin polypeptid s can be hybrid glutenin polypeptides comprising sequences from two or more different subunits or may be polypeptides in which single amino acid modifications are made.

In the case where an inserted polynucleotide sequence is transcribed and translated to produce a functional glutenin polypeptide, one of skill will recognize that because of codon degeneracy, a number of polynucleotide sequences will encod the same polypeptide. These variants are specifically cover d by the terms "qlutenin gene" or "glutenin polynucleotid In addition, the terms specifically include th se sequence". full length sequences substantially identical (determined as described below) with a glutenin gene sequence and that enc de proteins that retain the function of the glutenin polypeptide. Thus, in the case of wheat glutenin genes disclosed here, the term includes variant polynucleotide sequences which have substantial identity with the sequences disclosed here and which encode glutenin polypeptides capable of crosslinking by disulfide bonds with other glutenin polypeptides glutenin polymers.

Two polynucleotides or polypeptides are said to be "identical" if the sequence of nucleotides or amino acid residues, respectively, in the two sequences is the same who aligned for maximum correspondence as described below. The term "complementary to" is used herein to mean that the complementary sequence is identical to all or a portion of a reference polynucleotide sequence.

Sequence comparisons between two (or more) polynucleotides or polypeptides are typically performed by comparing sequences of the two sequences over a segment or "comparison window" to identify and compare local regions of sequence similarity. Optimal alignment of sequences for comparison may be conducted by the local homology algorithm of Smith and Wat rman Adv. Appl. Math. 2: 482 (1981), by the homol gy alignment algorithm of Needleman and Wunsch J. Mol. Bi l. 48:443 (1970), by the search for similarity method of P arson and Lipman Proc. Natl.

Sci. (U.S.A.) 85: 2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, G netics Comput r Group (GCG), 575 Science Dr., Madison, WI), or by "Percentage inspection. of sequence identity" determined by comparing two optimally aligned sequences ov r a comparison window, wherein the portion of the polynucleotide sequence in the comparison window may comprise additions or deletions (i.e., gaps) as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical nucleic acid base or amino acid residue occurs in sequences to yield the number of matched positions, dividing the number of matched positions by the total number positions in the window of comparison and multiplying the result by 100 to yield the percentage of sequence identity.

"substantial of identity" polynucleotide sequences means that a polynucleotide comprises a sequence that has at least 60% sequence identity, preferably at least 80%, more preferably at least 90% and most preferably at least 95%, compared to a reference sequence using the programs described above using standard parameters. One of skill will recogniz that these values can be appropriately adjusted to determine corresponding identity of proteins encoded by two nucleotide sequences by taking into account codon degeneracy, amino acid similarity, reading frame positioning and Substantial identity of amino acid sequences for these purposes normally means sequence identity of at least 40%, preferably at least 60%, more preferably at least 90%, and most preferably at least 95%. Polypeptides which are "substantially similar" share sequences as noted above except that residue positi ns which are not identical may differ by conservative amino acid changes. Conservative amino acid substitutions refer to the interchangeability of residues having similar side chains. example, a group of amino acids having aliphatic side chains is glycine, alanine, valine, leucine, and isol ucine; a group of amino acids having aliphatic-hydroxyl side chains is serine and threonin; a group of amin acids having amide-containing side chains is asparagine and glutamine; a group of amino acids

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having aromatic side chains is phenylalanin, tyrosine, and trypt phan; a group of amino acids having basic side chains is lysine, arginin, and histidine; and a group of amino acids having sulfur-containing side chains is cysteine and methionine. Preferred conservative amino acids substitution groups are: valine-leucine-isoleucine, phenylalanine-tyrosin, lysine-arginine, alanine-valine, and asparagine-glutamine.

indication that Another nucleotide sequences substantially identical is if two molecules hybridize to each other under stringent conditions. Stringent conditions are different and will be sequence dependent in different Generally, stringent conditions are selected to circumstances. be about 5°C to about 20°C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength and The Tm is the temperature (under defined ionic strength and pH) at which 50% of the target sequence hybridizes to a perfectly matched probe. However, nucleic acids which do not hybridize to each other under stringent conditions are still substantially identical if the polypeptides which they encode are substantially identical. This may occur, e.g., when a copy of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a diagram showing the domains of y-type and x-type HMW glutenins. The C- and N-terminal domains are shad d, the locations of cysteine residues are designated by S.

Figure 2A is a diagram showing the structure of a hybrid glutenin polypeptide of the invention.

Figure 2B is a diagram of the recombinant plasmid of the invention which encodes the polypeptide shown in Figure 2A.

Figure 3 is photograph of a gel stained with Coomassie blue showing the relative mobility of a hybrid HMW glutenin polypeptide of the invention (marked by the arrow).

DESCRIPTION OF THE PREFERRED EMBODIMENTS

This invention relates to plant glutenin genes, in particular wheat glutenin g nes. Nucl ic acid sequenc s from glutenin g n s can be used to modify glut nin content in transgenic plants, in particular wheat plants. In addition,

th inv ntion provides new seed-specific pr mot rs us ful for directing expr ssion of desired genes in a number of plants.

Generally, the nomenclature and the laboratory procedur s in recombinant DNA technology described b low are those well known and commonly employed in the art. Standard techniqu s are used for cloning, DNA and RNA isolation, amplification and purification. Generally enzymatic reactions involving DNA ligase, DNA polymerase, restriction endonucleases and the like are performed according to the manufacturer's specifications. These techniques and various other techniques are generally performed according to Sambrook et al., Molecular Cloning - A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, (1989).

Glutenin polypeptides and genes

The HMW glutenins are composed primarily of a central domain of 45-90 repeating two or three simple motifs, comprising chiefly glutamine and proline. This unusual primary structure results in a rod-like secondary structure as assessed by circular dichroism and SEM. Cysteines in the C- and N-terminal domains of the molecules are known to be critical fr the formation of intermolecular disulfide bonds (Figure 1). Th quaternary interactions of these proteins are of interest because the disulfide crosslinks among HMW and LMW subunits is thought to be integral to the elastic properties of dough.

Any isolated glutenin gene can be used in the present invention. The particular polynucleotide sequence used is not a critical feature of the invention, so long as the desired alteration in glutenin content is achieved. As noted above, HMW glutenins (both x-types and y-types) are encoded at th Glu-1 locus. In hexaploid wheat the group 1 chromosomes are designated 1A, 1B and 1D. Studies of the frequency of allel s at each of the loci indicate the presence of at least three alleles at Glu-Al, 11 alleles at Glu-Bl, and six alleles at Glu-Dl. The gene products of each allele are designated by noting the chromosome, followed by its classification as x ry, and a number (see, MacRitchi, Advancs in Food and Nutrition Research 36:1-87 (1992)).

What HMW glutenin g nes hav by n clond and described in the literature. Frinstanc, a complete set of HMW glutenin

gen s have been isolated and sequenced from the hard red winter wh at Cheyenne (Anderson t al. Wheat Genetics Symposium Proc. pp 699-704 (Bath Pr ss, Cambridge, UK, 1988), including two, Dx5 and Dy10, with th high st correlation with good flour quality (Anderson et al. Nuc. Acids. Res. 17:461-462 (1989)). Other particularly useful HMW glutenins include Ax and Bx alleles.

The isolation of other glutenin genes may be accomplished by a number of techniques. For instance, oligonucleotide probes based on the sequences disclosed in the prior art can be used to isolate the desired gene from a cDNA or genomic DNA library. To construct genomic libraries, large segments of genomic DNA are generated by random fragmentation, e.g. using restriction endonucleases, and are ligated with vector DNA to form concatemers that can be packaged into the appropriate vector. To prepare a cDNA library, mRNA is isolated from endosperm and a cDNA library which contains the glutenin gene transcript is prepared from the mRNA.

Alternatively, the nucleic acids of interest can samples using amplification amplified from nucleic acid techniques. For instance, polymerase chain reaction (PCR) technology to amplify the sequences of the glutenin and relat d genes directly from genomic DNA, from cDNA, from genomic libraries or cDNA libraries. PCR and other in vitro amplification methods may also be useful, for example, to clon nucleic acid sequences that code for proteins to be expressed, to make nucleic acids to use as probes for detecting the presence of the desired mRNA in samples, for nucleic acid sequencing, or for other purposes. For a general overview of PCR see PCR Protocols: A Guide to Methods and Applications. (Innis, M, Gelfand, D., Sninsky, J. and White, T., eds., Academic Press, San Diego (1990)).

Polynucleotides may also be synthesized by well-known techniques as described in the technical literature. See, e.g., Carruthers et al., Cold Spring Harbor Symp. Quant. Biol. 47:411-418 (1982), and Adams et al., J. Am. Chem. Soc. 105:661 (1983). Double stranded DNA fragm nts may then be obtain d either by synthesizing the complementary strand and ann aling the strands t gether under appropriate conditions, or by adding

th complem ntary strand using DNA polymerase with an appropriate prim r sequence.

Isolated s quences prepared as describ d herein can then bound to modify glutening ne expression and therefor glutenin content in plants. One of skill will recognize that the nucleic acid encoding a functional glutenin protein need not have a sequence identical to the exemplified general school of the exemplified general polypeptides can be used in the present invention.

noted above, glutenin polypeptides, like other have different domains which perform different functions. Thus, the glutenin gene sequences need not be full length, so long as the desired functional domains of the protein is expressed. Chimeric glutenin polypeptides can be readily designed utilizing various recombinant DNA techniqu s well known to those skilled in the art. For example, the chains can vary from the naturally occurring sequence at the primary structure level by amino acid substitutions, additions, deletions, and the like. In particular, cysteine residues may be added, deleted or moved within the polypeptide to achieve a modified glutenin polypeptide with desired properties. Chimeric polypeptides may also be produced by fusing coding sequences from two or more glutenin genes. All of these modifications can be used in a number of combinations to produce the final modified protein chain.

Preparation of recombinant constructs

To use isolated glutenin sequences in modifying glutenin in plants, recombinant DNA vectors suitable transformation of plant cells are prepared. A DNA sequence coding for the desired glutenin polypeptide, for example a cDNA or a genomic sequence encoding a full length protein, construct a recombinant conveniently used to expression cassette which can be introduced into the desired plant. expression cassette will typically comprise the glut nin polynucleotide sequence operably linked to a promoter sequ nce other transcriptional and translational r gulatory sequ nces which will dir ct the transcription of the s quence from the glutenin g ne in the int nd d tissues (.g., ndosperm) of th transformed plant.

For example, a constitutive plant prom ter fragment may be employ d which will direct expression of th glutenin in all Such promoters are active under most tissues of a plant. envir nm ntal conditions and states of developm nt or cell differentiation. Examples of constitutive promoters includ cauliflower mosaic virus (CaMV) 35S transcription the initiation region, the 1'- or 2'- promoter derived from T-DNA tumafaciens, Agrobacterium and other transcription initiation regions from various plant genes known to those f skill.

Alternatively, the plant promoter may be under environmental control. Such promoters are referred to here as "inducible" promoters. Examples of environmental conditions that may effect transcription by inducible promoters include pathogen attack, anaerobic conditions, or the presence of light.

Typically, the promoters used in the constructs of the invention will be "tissue-specific" and are under developmental control such that the desired gene is expressed only in certain tissues, such as leaves, roots, fruit, seeds, or flowers. Promoters that direct expression in seeds, particularly the endosperm are particularly preferred. Examples of such promoters include the promoter from genes encoding seed storage proteins, such as napin, cruciferin, phaseolin, and the like (see, U.S. Patent No. 5,420,034). Other promoters suitable for expressing glutenin genes in cereals include promoters from genes encoding gliadins, cereal prolamines (e.g., zein, hordein, secalin, and avenin) and starch biosynthetic enzymes.

endogenous promoters from glutenin genes The particularly useful for directing expression of glutenin genes to the seed, particularly the endosperm. These seed-specific promoters can also be used to direct expression of heterologous structural genes. Thus, the promoters can be used recombinant expression cassettes to drive expression of any gene whose expression in seeds is desirable. Examples include genes encoding proteins useful in increasing the nutritional value of seeds (e.g., gen s ncoding proteins involved in lipid, protein, and carbohydrate or starch biosynthesis). Oth r g n s include those encoding pharmaceutically us ful

compounds, and genes no ding plant r sistance products to combat fungal or other inf ctions of th seed.

The glut nin promoters can also b us d to initiate transcription of mRNA molecules t inhibit expression of endogenous endosperm gene. Exemplary genes whose expression could be inhibited include genes encoding glutenins, enzymes involved in carbohydrate or lipid synthesis, secalins, and th Means for inhibiting gene expression in plants using recombinant DNA techniques are well known. For instance, antisense technology can be conveniently used. Sheehy et al., Proc. Nat. Acad. Sci. USA, 85:8805-8809 (1988), and Hiatt et al., U.S. Patent No. 4,801,340. Catalytic RNA molecules or ribozymes can also be used to inhibit expression of endosperm-specific genes. The design and use of target RNAspecific ribozymes is described in Haseloff et al. Nature, 334:585-591 (1988). Introduction of nucleic acid configured in the sense orientation has also been shown to be an effective means by which to block the transcription of target genes. an example of the use of sense suppression to modulate expression of endogenous genes see, Napoli et al., The Plant 2:279-289 (1990), and U.S. Patents Nos. 5,034,323, 5,231,020, and 5,283,184.

The glutenin promoters of the invention are typically at least about 400 base pairs in length, and often at least about 800 or about 1000 base pairs. The length of the promoters is typically less than about 3500 base pairs, usually less than about 2800 base pairs and often less than about 2000 base pairs in length. The length of the promoters is counted upstream from the translation start codon of the native gene. One of skill will recognize that use of the "about" to refer to lengths of nucleic acid fragments is meant to include fragments of various lengths that do not vary significantly from the lengths recited here and still maintain the functions of the claimed promoters (i.e., seed-specific gene expression).

To identify glutenin promoters, the 5' portions of a genomic glutenin gene clone is analyzed for sequences characteristic of promot r s qu nces. For instance, promot r s quence elements includ the TATA box consensus sequence (TATAAT), which is usually 20 to 30 bas pairs upstr am of th transcription start site. In plants, further upstream from th

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TATA box, at positions -80 to -100, there is typically a promoter element with a series of adenines surrounding the trinucleotide G (or T) N G. J. Messing et al., in Genetic Engineering in Plants, pp. 221-227 (Kosage, Meredith and Hollaender, eds. 1983).

In preparing expression vectors of the invention, sequences other than the promoter and the structural gene of interest are also preferably used. If proper polypeptid expression is desired, a polyadenylation region at the 3'-end of the glutenin coding region should be included. The polyadenylation region can be derived from the natural gene, from a variety of other plant genes, or from T-DNA.

The vector comprising the sequences from a glutenin gene will typically comprise a marker gene which confers a selectable phenotype on plant cells. For example, the mark r may encode biocide resistance, particularly antibiotic resistance, such as resistance to kanamycin, G418, bleomycin, hygromycin, or herbicide resistance, such as resistance to chlorosluforon, or phosphinothricin (the active ingredient in bialaphos and Basta).

Preparation of transgenic plants

The DNA constructs described above may be introduced into the genome of the desired plant host by a variety of conventional techniques. Techniques for transforming a wid variety of higher plant species are well known and described in the technical and scientific literature. See, for example, Weising et al., Ann. Rev. Genet. 22:421-477 (1988).

The DNA construct may be introduced directly into the genomic DNA of the plant cell using techniques such as biollistic methods, electroporation, PEG poration, and microinjection of plant cell protoplasts or embryogenic callus. Alternatively, the DNA constructs may be combined with suitable T-DNA flanking regions and introduced using an Agrobacterium tumefaciens or A. rhizogenes vector.

Particle bombardment techniques are described in Klein t al., Nature 327:70-73 (1987). A particularly preferred method of transforming wheat and oth r cer als is the bombardment of calli derived from immature embryos as described by Weeks t al. Plant Physi 1. 102:1077-1084 (1993).

Microinj ction t chniques are known in the art and well describ d in the scientific and patent literature. The introduction of DNA constructs using polythylen glycol precipitation is described in Paszkowski t al. Embo J. 3:2717-2722 (1984). Electroporation techniques are described in Fromm et al. Proc. Natl. Acad. Sci. USA 82:5824 (1985).

tumefaciens-meditated Agrobacterium transformation techniques are also well described in the literature. See, for example Horsch et al. Science 233:496-498 (1984), and Fraley et al. Proc. Natl. Acad. Sci. USA 80:4803 (1983). Although Agrobacterium is useful primarily in dicots, certain monocots can be transformed by Agrobacterium. instance, Agrobacterium transformation of rice is described by Hiei et al. Plant J. 6:271-282 (1994).

The present invention is particularly useful in wheat and other cereals. A number of methods of transforming cereals been described in the literature. For instance, described by transformation of rice is Toriyama Bio/Technology 6:1072-1074 (1988), Zhang et al. Theor. Appl. Gen. 76:835-840 (1988), and Shimamoto et al. Nature 338:274-276 Transgenic maize regenerants have been described by Fromm et al., Bio/Technology 8:833-839 (1990) and Gordon-Kamm et al., Plant Cell 2:603-618 (1990)). Similarly, oats (Sommers et al., Bio/Technology 10:1589-1594 (1992)), wheat (Vasil et al., Bio/Technology 10:667-674 (1992)); Weeks et al., Plant Physiol. 102:1077-1084 (1993)), sorghum (Casas et al., Proc. Natl. Acad. Sci. USA 90:11212-11216 (1993)), rice (Li et al., Plant Cell Rep. 12:250-255 (1993)), barley (Yuechun and Lemaux, Plant Physiol. 104:37-48 (1994)), and rye (Castillo et al., Bio/Technology 12:1366-1371 (1994)) have been transformed via bombardment.

Transformed plant cells which are derived by any of the above transformation techniques can be cultured to regenerate a whole plant which possesses the transformed genotype and thus the desired phenotype. Such regeneration techniques rely on manipulation of certain phytohormones in a tissue culture growth medium, typically r lying on a biocid and/or h rbicid mark r which has been introduced together with the glut nin polynucl otide s quences. Plant regeneration from cultured protoplasts is described in Evans t al., Pr toplasts Isolation

and Cultur, Handbook of Plant Cell Culture, pp. 124-176, MacMillilan Publishing Company, New York, 1983; and Binding, Reg neration of Plants, Plant Protoplasts, pp. 21-73, CRC Press, Boca Raton, 1985. Regeneration can also be btained from plant callus, explants, organs, or parts thereof. Such regeneration techniques are described generally in Klee et al. Ann. Rev. of Plant Phys. 38:467-486 (1987).

The methods of the present invention are particularly useful for incorporating the glutenin polynucleotides into transformed plants in ways and under circumstances which are not found naturally. In particular, the glutenin polypeptides may be expressed at times or in quantities which are not characteristic of natural plants.

One of skill will recognize that after the expression cassette is stably incorporated in transgenic plants and confirmed to be operable, it can be introduced into other plants by sexual crossing. Any of a number of standard breeding techniques can be used, depending upon the species to be crossed.

The invention has use over a broad range of types f plants. The glutenin genes are preferably expressed in cereal species commonly used for production of flour, e.g., wheat, rye, oats, and the like.

The seed-specific promoters from glutenin genes can be used in essentially any plant species. For instance, the promoters or genes described here can be used in species from genera Asparagus, Atropa, Avena, Brassica, Citrullus, Capsicum, Cucumis, Cucurbita, Daucus, Fragaria, Glycine, Gossypium, Helianthus, Heterocallis, Hyoscyamus, Juglans, Lactuca, Linum, Lolium, Lycopersicon, Malus, Manihot, Majorana, Medicago, Nicotiana, Oryza, Panieum, Pannesetum, Persea, Phaseolus, Pisum, Pyrus, Prunus, Raphanus, Secale, Senecio, Sinapis, Solanum, Sorghum, Triticum, Vitis, Vigna, and, Zea. Plants that are particularly useful in the invention include Avena, Brassica, Glycine, Hordeum, Oryza, Phaseolus, Pisum, Secale, Sorghum, Triticum, Vigna, and Zea.

As not d above, the inv ntion is particularly useful for improvement of wheat cultivars. Any wheat cultivar may be improved by the methods of th invention. Exemplary lines

include hexaploid lin s such as Anza, Shasta, Y coro R jo, Siouxland, Freedom, Tam107, and Tam200.

The effect of the modification of glutenin g ne expr ssion can be measured by detection f incr as s or decreas s in desired glutenin protein levels using, for instance, gel electrophoresis, as described below. Quantification of HMW glutenin content can be carried out by SDS/PAGE densitometry, as described below.

Other methods for quantifying glutenin content include sonication of flour dispersions in SDS buffer to solubilize glutenin and other relatively insoluble proteins (see, Singh and MacRitchie in Wheat End-Use Properties: Wheat and Flour Characterization for Specific End-Uses H. Salovaara, ed., pp 321-326 (University of Helsinki, Helsinki, 1989). This method is useful in solubilizing at least 95% of total flour protein. Quantification of the glutenin fraction can then be performed using size exclusion high performance liquid chromatography as described by Batey et al., Cereal Chem. 68:207-209 (1991). Other suitable methods include reverse phase HPLC and capillary electrophoresis.

The plants of the invention have increased in glutenin content in the endosperm of mature seeds as compared to the parental lines from which they are derived. The increase is usually at least about 15% as compared to the parental line. More usually, the increase is at least about 20%, preferably at least about 35%, and in more preferred embodiments at least about 45%. Any of the methods for quantifying glutenins in the endosperm described above can be used to determine the percent increase in glutenin content. A wheat seed typically reaches maturity about 6 to about 10 weeks after anthesis.

Another measure of increased glutenin content in transgenic plants of the invention is glutenin content as a percent of total protein of a mature seed or flour derived from the mature seed. Different wheat cultivars have different glutenin contents as measured using standard methods. Thus, the final glutenin content of the plants of the invention will d pend upon th parental line. In the case where HMW glutenin genes ar used, the plants of th invention will typically hav HMW glut nin contents f at least ab ut 15%, usually at least

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about 18%, and preferably at least about 20% as measur d using standard methods such as thos describ d in detail b low.

Analysis of th rh ological properties of flour deriv d from the transgenic plants of the invention can be carried out according to standard physical dough-testing instruments widely used to measure flour and dough quality (see, e.g., MacRitchie, Advances in Food and Nutrition Research 36:1-87 (1992)). methods include, for instance, use of extensographs to measure tensile strength. Two of the main parameters measured are maximum resistance (Rmx) and extensibility (Ext). Other mixographs methods include and bake-test loaf volume (MacRitchie and Gras Cereal Chem. 50:292-302 (1973) as well as SDS-sedimentation tests, amelographs, and cookie methods.

The following Examples are offered by way of illustration, not limitation.

Example 1

This example demonstrates increase in glutenin content in transgenic wheat plants comprising recombinant constructs of the invention.

MATERIALS AND METHODS

Construction of the hybrid HMW glutenin expression plasmid

Two HMW glutenin subunits, Dx5 and Dy10, encoded by the Glu-Dl-lb and Glu-Dl-2b genes, respectively, have been most often associated with superior dough strength by genetic correlation studies. A bacterial expression vector comprising a fusion of coding regions of the Dy10 and Dx5 genes is described by Shani et al., Plant Physiol. 98:433-441 (1992). The hybrid HMW glutenin polypeptide encoded by this gene is diagrammed in Figure 2A.

In this example, the Dy10:Dx5 hybrid coding region was reunited with 5' and 3' flanking sequences from the native Dy10 and Dx5 genes. The construction of a fusion at the HindIII site located at base pairs 429-435 and 385-391 after the A of the start cod ns of the Glu-Dl-2b (Genbank Accession No. X12929) and Glu-Dl-1b (G nbank Accession No. X12928) genes (And rson et al. Nuc. Acids Res. 17:461-462 (1989)) and its

insertion into the bacterial expression v ctor pet3A to form p t-3a-10/5 is described by Shani et al., supra.

th native Dyl0 gene provid s th transcribed untranslat d 5 ' transcription start sit, th region and the first 145 codons including the 21 amino acid signal peptide. The native Dx5 gene provides its codons 130 to 848 followed by two translation stop codons, the transcription termination and poly(A) addition signals. Because amino acids 138 through 147 are shared by the Dy10 and Dx5 glutenin subunits, the central region of repeating amino acid motifs of the hybrid glutenin polypeptide (thickest boxes in Figure 2A) identical to that of Dx5. The configuration of th cysteines (S) and thus of potential disulfide bonds is chimeric in nature: the N-terminal cysteines are of the y-type and the single C-terminal cysteine is typical of x-type subunits.

reconstitute the promoter In order to and any transcription start and termination sites, regulatory signals needed for expression in wheat, fragments were isolated and combined in a single ligation reaction with the Bluescript KS (Stratagene) vector cut with 1) the 715 bp StuI/DrdI fragment that EcoRI and BamHI: contains the junction region in pET-3a-10/5; 2) the 2800 bp fragment from a clone of the native Dy10 gene beginning at the EcoRI site 5' to the gene and ending at the StuI site 74 bp after the A in the start codon; and 3) the 3800 bp fragment from a clone of the native Dx5 gene that starts at the DrdI site 744 bp after the A in the start codon and ends at a BamHI site in the vector just outside the EcoRI site in the 3' A plasmid with the correct structure was flanking region. identified by restriction analysis and named pGlu10H5 (Figur This plasmid was deposited under the terms of the Budapest Treaty in the Agricultural Research Culture Collection (NRRL), 1815 North University Street, Peoria, Illinois, USA, on January 12, 1996 and has been assigned Accession No. NRRL B-21517.

The plasmid UBI:BAR which comprises the maize ubiquitin promoter operably link d to the bar gene ncoding the nzyme PAT, which inactivats phosphinothricin was prepard as described Cornejo et al. Plant Mol. Biol. 23:567-581 (1993).

Both plasmids w re prepared by the alkaline lysis meth d using a Qiagen kit for the final purificati n. The DNAs were stored in TE buffer at a conc ntration of 1 mg/ml.

Wheat Transformation

12.5 ug of each DNA (an approximately 2:1 molar ratio f UBI:BAR to pGlul0B5) was coated onto gold particles for bombardment into immature wheat embryos of cultivar Bobwhite essentially as described by Weeks et al. Plant Physiol. 102:1077-1084 (1993) except that the embryos were incubated on callus induction media containing mannitol for 4 hours before and 20 hours after bombardment. Transformants were selected on the basis of their resistance to lmg/L bialaphos at the callus and green shoot stages of regeneration and to 3 mg/L bialaphos at the rooting stage of regeneration as described by Weeks et al., supra.

Protein Analysis

Protein extracts from immature wheat endosperm wer prepared by grinding the tissue or cells in SDS-PAGE sample loading buffer [66 mM Tris pH 6.8, 3% (w/v) SDS, 0.05% (w/v) Bromphenol blue, 2% (v/v) b-mercaptoethanol]. Protein extracts from mature dry wheat seeds were prepared by dissolving flour in sample buffer. SDS-PAGE was according to standard procedures. The separation gels for detection and quantitation of the hybrid subunit were 10% (w/v) acrylamide 0.05% (w/v) bis-acrylamide. The gels were run at 40 mamps constant current until 20 minutes after the bromphenol blue dye front had run off the gel (about 1000 Volt-hours).

The gels were stained by the Neuhoff method with a colloidal suspension of Coomassie blue. Protein standards for molecular weight calibration were purchased from Pharmacia and consisted of Phosphorylase b (94 kDa) Bovine Serum Albumin (67 kDa), Ovalbumin (43 kDa), Carbonic Anhydrase (30 kDa), Soybean Trypsin Inhibitor (20.1 kDa) and a-Lactalbumin (14.4 kDa). Stained gels were scanned using an Alpha Innotech densitometer.

DNA Analysis

Southern analyses of genomic DNA was as describ d in Weeks t al., supra, except that 10 ug of DNA w re loaded into each

slot of a 0.6% agarose gel. The pr b consisted of th 1578 bp PvuII fragment from the coding r gion of Dx5 (bracketed in Figur 2B).

RESULTS

Forty-five independent callus pieces were selected from 4980 embryos bombarded in 10 different experiments; each yielded one to five plants. The To primary regenerated plants and the lines derived from them are designated with an unique number following the bombardment number, eg., plant 126 regenerated from bombardment 7 is 7-126. Plants from twenty-six of these independent line gave rise to progeny embryos that were able to germinate in the presence of 3 mg/L bialaphos when excised from immature seeds three to four weeks after anthesis.

The immature endosperm tissue corresponding to each excised embryo was extracted and screened by SDS-PAGE for th presence of the hybrid glutenin. This SDS-PAGE analysis showed that the migration of the hybrid HMW glutenin subunit is clearly distinguishable from the native HMW subunits und r these gel conditions.

Although the lanes did not contain the same amounts f total protein, it was evident that there is variation in the expression levels of the hybrid subunit relative to those of the native HMW subunits in the same seed. Of the twenty-six resistant lines that were derived from co-bombardm nt experiments including pGlulOH5, sixteen (60%) lines show d expression of the hybrid subunit in the T, generation.

Expression stability

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At each successive generation, 24 or 25 immature embryos 3-4 weeks after anthesis were tested for their ability to germinate on bialaphos. Plants were classified as homozygous if all 25 of their progeny embryos exhibited resistance. levels of transgene expression were assessed in successive generations. In most cases, expression was stable. An increase in expression between the T1 T2 generations in some plants was explain d by the incr as gene dosage between th h terozygous and homozygous progeny of th T0 plant and T1 generation, respectively. the endosperm is triplicate, individual selfed prog ny of th To

heter zygot could have 0, 1, 2 or 3 copi s and th T1 lane contains an average of 8 of thes individuals.

In four of th 15 wheat lines analyzed, expr ssion of the hybrid HMW glut nin has not remained at the same levels, but rather has declined as homozygous resistant lines were selected in the T₂ or the T₃ generations. The decline could be du either to hybrid HMW glutenin transgene inactivation or to loss of unlinked pGlu10H5 transgene(s) copies via segregation.

Cross-linking ability of the hybrid subunit

As noted above, the hybrid HMW glutenin contains a uniqu configuration of cysteines, y-type at the N-terminus and x-type at the C-terminus. The hybrid HMW glutenin behaved exactly like the native subunits in terms of its solubility in SDS buffers with and without a reducing agent, when examined by SDS-PAGE. The native and hybrid HMW glutenins were partially extracted from both the Bobwhite and the transgenic flours by SDS alone and were present in lower molecular weight polymers. The majority of the hybrid glutenin, however, was insoluble in SDS buffer and could only be extracted when a reducing agent was added. Thus, the configuration of cysteines in the hybrid glutenin does not preclude the formation of intermolecular disulfide bonds or high molecular weight polymers.

Relative expression levels

Figure 3 shows SDS-PAGE analysis of protein extracts from eight different transgenic lines (Lanes 1-8) and Bobwhite (C). Each sample was prepared from flour of a mixture of eight mature seeds of a single homozygous T₂ or T₃ plant. Under the segliconditions, the native By9 and Dy10 subunits are not well resolved and run as a closely spaced doublet of apparent molecular weight 94 kDa. Likewise the two isoforms of the Dx5 subunit are not well-resolved and migrate as a broad band. The Bx7 band is well-separated from other proteins and the darkest of the HMW subunits in intensity. In all the lines except the one shown in Lane 3, the hybrid HMW subunit is pr s nt in at least as high a quantity as the native Ax2* (larg st) HMW subunit. Thus even a single copy of the hybrid transgen (Lane

1) can support significant expression 1 vels, comparabl to those of the native HMW subunits.

In on of the lines (shown in lane 2), synthesis of native HMW glutenin synth sis has declined relative to the other proteins. The extract shown there exhibits reduced synthesis of all the native HMW glutenin compared to the lower molecular weight seed proteins in the same lane.

In order to estimate the levels of the hybrid HMW glutenin relative to those of the native HMW glutenin and other s ed proteins, the gel was scanned by densitometry. All the wh at prolamines stay on the gel. However, since proteins f molecular weights less than 33 kDa are run off the gel under the conditions used to separate the hybrid HMW glutenin from the nearby Dx5 subunit band, only relative quantities could be assessed. The results of these comparisons are shown in Tabl I along with the number of intact transgene copies estimated from restriction enzyme digests of genomic DNA (see below). The hybrid HMW glutenin band constitutes more than 20% of the total HMW glutenins in all the lines except that in lane 3. Thus the transgene has made a significant contribution to the HMW glutenin composition in all the transgenic lines.

In the case where endogenous HMW glutenin accumulation is reduced (Lane 2), the hybrid makes up 62% of the total complement of HMW glutenin. In most lines, the sum of the HMW glutenin accumulation is elevated 29 to 45% relative to the HMW glutenin content of Bobwhite (all sums are normalized to the remainder of the protein their respective lanes). Thus this quantitative analysis suggests that addition of HMW-glutenin gene copies raises the levels of HMW glutenin accumulation in mature seeds.

DNA analysis

DNA was isolated from the leaves of transgenic plants of each line and assayed for the presence of transgenes by genomic Southern analysis, as described above. Hybridizing bands known to correspond to the transgenes were scanned and their d nsities compared. The results are shown in Table I. The numb r ranges from 1 in line 11-291 to about 6 in 14-23 (Table I). This range is much br ader than that in total HMW glutenin levels or even hybrid HMW glutenin levels. However the most

highly expressing lin had one of th highest copy numbers (Line 4) and th poor st expr ssion was in a single copy line (Line 3).

Table I. Transgenic HMW-glutenin gene activity in individual wheat lines.

Line	Transgene Expression	Total H subunit as % total	MW Gene Copy No. of Estimate
1	23%	135%	1
2	62%	70%	5-6
3	15%	145%	1
4	42%	145%	4-5
5	23%	140%	2
6	24%	129%	2
7	22%	135%	3-4
8	20%	143%	3-4

DISCUSSION

The experiments reported here show that levels of HMW glutenins can be increased by addition of gene copies. The behavior of the hybrid subunit is indistinguishable from that of the native subunits. The protein accumulates in the endosperm tissue over the course of seed development and is cross-linked by disulfide bonds into large polymers that cannot be extracted by SDS without addition of reducing agents.

Expression of the hybrid glutenin is under the control of the native DylO 5' and Dx5 3' flanking sequences. These sequences were sufficient to achieve high levels of the modified seed storage protein. Even a single gene copy supported expr ssion levels comparable to thos of th nativ HMW glutenin g nes. Thus, th se s qu nces serv as a source f effective regulatory s quences suitable f r expression of oth r

proteins in transgenic wheat and potentially other cereal endosperm.

Example 2

This example describes the transformation of wheat using a recombinant construct encoding Dx5 and Dy10.

The recombinant expression vectors were prepared as generally as described above except that the complete Dx5 or Dy10 gene, including 5' and 3' regulatory sequences, were used. The constructs were then introduced into immature wheat embryos by particle bombardment as described above. Analysis of T_0 plants and their progeny is performed as described above.

The above examples are provided to illustrate the invention but not to limit its scope. Other variants of the invention will be readily apparent to one of ordinary skill in the art and are encompassed by the appended claims. All publications, patents, and patent applications cited herein are hereby incorporated by reference for all purposes.

Applicant's or agent's file	International application Mo
reference number A700/363	

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WHAT IS CLAIMED IS:

1. A method f altering glutenin c ntent in the ndosperm of a wheat plant, the meth d comprising:

stably transforming a parental wheat plant with a recombinant expression cassette comprising a nucleic acid encoding a glutenin polypeptide and a seed-specific promoter to produce a stably transformed wheat plant; and

selecting a progeny of said stably transformed wheat plant having a glutenin content in the endosperm of a mature seed at least about 30% different than the glutenin content in the endosperm of a mature seed from the parental wheat plant.

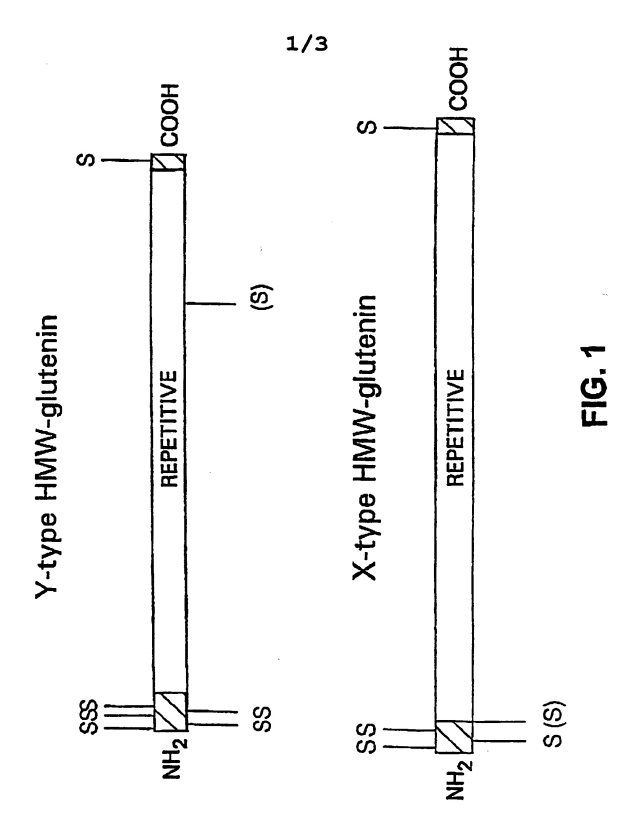
- 2. The method of claim 1, wherein the nucleic acid is stably transformed into the first wheat plant using particl bombardment.
- 3. The method of claim 1, wherein the nucleic acid encodes a chimeric glutenin polypeptide.
- 4. The method of claim 1, wherein the chimeric polypeptide comprises sequences from an x-type glutenin polypeptide and a y-type glutenin polypeptide.
- 5. The method of claim 1, wherein the x-type polypeptide is encoded by the Glu-Dl-lb gene.
- 6. The method of claim 1, wherein the y-type polypeptide is encoded by the Glu-D1-2b gene.
- 7. The method of claim 1, wherein the promoter is from the Glu-Dl-2b gene.
- 8. The method of claim 1, wherein the parental wheat plant is Bobwhite.
- 9. A wheat plant comprising a recombinant expr ssion cass tte comprising a nucl ic acid nc ding a glut nin polypeptide and a s ed-specific promoter, the plant having a

glutenin content in the endosperm of a matur seed at least about 30% different than the glutenin cont nt in the endosperm of a mature seed from a parental wheat plant.

- 10. The plant of claim 9, wherein the nucleic acid encodes a chimeric glutenin polypeptide.
- 11. The plant of claim 10, wherein the chimeric polypeptide comprises sequences from an x-type polypeptide and a y-type glutenin polypeptide.
- 12. The plant of claim 11, wherein the x-type polypeptide is encoded by the Glu-D1-1b gene.
- 13. The plant of claim 11, wherein the y-type polypeptid is encoded by the Glu-D1-2b gene.
- 14. The plant of claim 9, wherein the promoter is from the Glu-Dl-2b gene.
- 15. The plant of claim 9, wherein the high molecular weight (HMW) glutenin accounts for at least about 15% of th total protein in the endosperm of the mature seed.
- 16. The plant of claim 9, wherein the parental wheat plant is Bobwhite.
 - 17. A seed derived from the plant of claim 9.
- 18. A recombinant construct comprising a wheat glutenin gene promoter of about 400 to about 2800 nucleotides operably linked to a heterologous polynucleotide sequence encoding a chimeric glutenin polypeptide, wherein the chimeric glutenin polypeptide is encoded by polynucleotide sequences from th Glu-Dl-lb gene and the Glu-Dl-2b gene.
- 19. The recombinant construct of claim 18, wh r in the wh at glutenin gene is Glu-D1-2b.

20. A plant comprising the recombinant construct of claim 18.

21. Wheat flour derived from the plant of claim 1.



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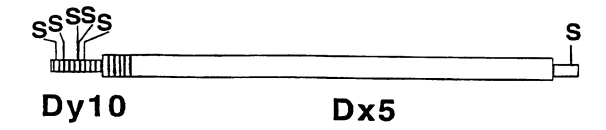


FIG. 2A

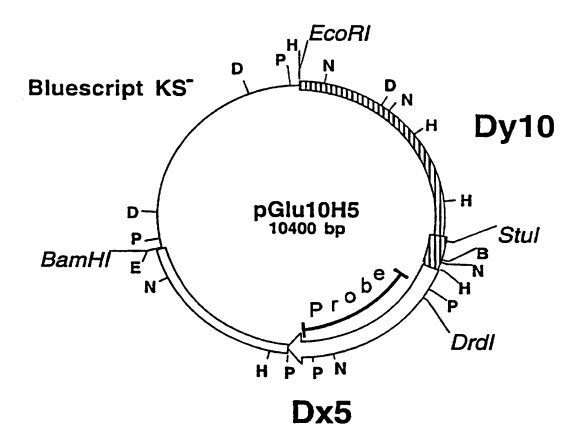


FIG. 2B

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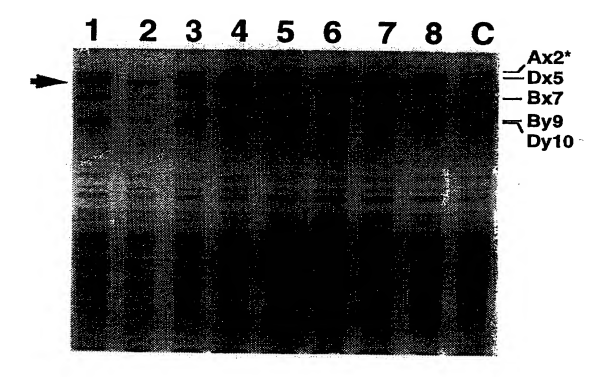


FIG. 3

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/00510

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APS, CA	BA, MEDLINE, CAPLUS, GENBANK, BIOSIS erms: wheat, glutenin, transformation, Glu-Di		where practicable	, search terms used)
C. DOC	UMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where	appropriate, of the rele	vant passages	Relevant to claim No.
Y	SHANI et al. Role of amino- and of the folding and oligomerization weight glutenin subunits. Plan pages 433-441, see entire document	of wheat high t Physiol. 1992	molecular	1-21
Y	WEEKS et al. Rapid production of fertile transgenic wheat (7) Physiol. 1993, Vol. 102, pages 101078-1079.	1-21		
Υ	ANDERSON et al. Nucleotide se molecular-weight glutenin genes hexaploid bread wheat, <i>Triticum</i> Nucleic Acids Research. 1989, V 462, see entire document.	from the D-ge	nome of a Cheyenne.	1-21
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INTERNATIONAL SEARCH REPORT

International application No.
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Categ ry*	Citation f document, with indicati n, where appropriate, of the relevant passages	Relevant to claim No
,	GALILI, G. Heterologous expression of a wheat high molecular weight glutenin gene in <i>Escherichia coli</i> . Proc. Natl. Acad. Sci. USA. October 1989, Vol. 86, pages 7756-7760, see entire document.	1-21
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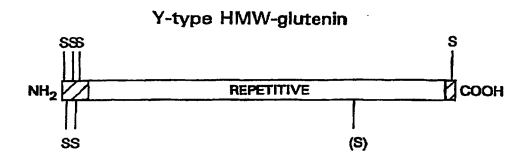
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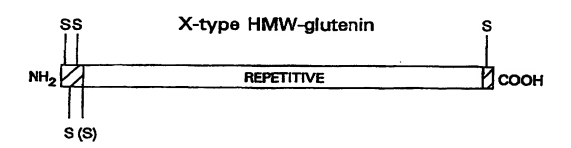
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Published

With international search report.

(54) Title: GLUTENIN GENES AND THEIR USES





(57) Abstract

The present invention is directed to methods of improving flour by altering glutenin content in seeds of plants, particularly wheat. The invention relates to methods of introducing a recombinant c nstruct comprising a glutenin gene into a parental plant.

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CS	Czechoslovakia	LT	Lithuania	TD	Chad
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GLUTENIN GENES AND THEIR USES

Field of the Invention

The present invention relates generally to plant molecular biology. In particular, it relates to nucleic acids and methods for improving glutenin content of plants.

BACKGROUND OF THE INVENTION

The glutenins, which include both high molecular weight (HMW) glutenin subunits and low molecular weight (LMW) glutenin subunits, comprise an economically important class of wheat seed storage proteins. The apparent molecular weights of the individual HMW glutenin polypeptides or subunits range from 90 to 200 kDa. These subunits crosslink by disulfide bonds among themselves and with LMW glutenin polypeptides to form polymers exceeding one million daltons in molecular weight. HMW glutenins constitute 8-10%, while LMW glutenins constitute 15-20% of the total endosperm protein. Both HMW and LMW glutenin proteins play important functional roles in determining the end-uses of wheat flour.

In wheat, HMW glutenins are encoded at the Glu-1 loci on the long arms of the group 1 chromosomes. Each locus consists of two separate genes, encoding an x-type and a y-type subunit, respectively. These pairs have never been confirmed to be separated by recombination. This has made determination of their separate contributions to bread dough properties difficult to assess by genetic correlation studies. For a review of the genetics and biochemistry of glutenin polypeptides, see, Shewry et al., J. Cereal Sci. 15:105-120 (1992).

Both the quantity and identity of specific HMW glutenin alleles contribute to the differences in bread-making quality of various cultivars. For instance, deletion of glutenin g n s results in a decrease in the overall levels of HMW glutenins, which results in decreases in bread-making quality (see, e.g., Lawrence et al. J. Cereal Sci 7:109-112 (1988)).

The effects of verproducing HMW glut nin on protein accumulation and baking quality has not been assess d becaus such lines of wheat have not been found among natural

populations. In addition, direct alteration of the glutenin subunits that f rm the polymers is not possibl using standard br eding methods. Thus, the art lacks repr ducible and efficient methods of producing lines with altered glutenin contents. The present invention addresses these and other needs.

SUMMARY OF THE INVENTION

The present invention provides methods of increasing glutenin in the endosperm of wheat plants. The methods and plants of the invention therefore are useful in providing flour and dough having improved end-use properties. The methods comprise introducing into a parental wheat plant a recombinant expression cassette comprising a nucleic acid encoding a glutenin polypeptide and selecting progeny wheat plant having increased glutenin content in the endosperm of mature seed. The glutenin content of the progeny is preferably at least about 15% greater than the glutenin of the parental wheat plant.

Any method of introducing the expression cassette into the parental plant can be used. Particle bombardment is a convenient method for producing transgenic plants. Once the expression cassette is stably integrated into the genom, standard sexual crosses can be used to introduce the expression cassette into desired lines.

In some embodiments, the nucleic acid introduced into the plant encodes a chimeric glutenin polypeptide. For instance, the chimeric polypeptide may comprise sequences from an x-typ glutenin polypeptide and a y-type glutenin polypeptide. Exemplary genes for this purpose include the Glu-Dl-lb gene and the Glu-Dl-2b gene.

The expression cassette may further comprise a seed-specific promoter to direct expression of the introduc d nucleic acid to the endosperm. A convenient promoter for this purpose is the promoter from the Glu-D1-2b gene.

Any wheat cultivar can be used as the parental line in the present invention. An exemplary cultivar is Bobwhite.

The inv nti n also provides wheat plants comprising a recombinant xpression cassett comprising a nucleic acid encoding a glut nin polyp ptide. The plants of the invention

have a glutenin cont nt in th endosp rm of a mature seed at least about 15% gr ater than the glutenin content in th endosperm of a mature seed from a par ntal wheat plant. The percentag of total endosperm protein which is glutenin will usually depend upon the glutenin content of the parental line. Usually, HMW glutenins account for at least about 15% of th total protein in the endosperm of the mature seed from plants of the invention.

The invention further provides recombinant constructs comprising a wheat glutenin gene promoter of about 400 to about 2800 nucleotides operably linked to a heterolog us polynucleotide sequence. These constructs are particularly useful in directing expression of the heterologous sequence to seeds of transgenic plants. An exemplary wheat glutenin gene from which the promoters of the invention can be derived is Glu-Dl-2b.

Definitions

The term "plant" includes whole plants, plant organs (e.g., leaves, stems, roots, flowers, etc.), seeds and plant cells and progeny of same. The class of plants which can bused in the methods of the invention is generally as broad as the class of higher plants amenable to transformation techniques, including both monocotyledonous and dicotyledonous plants.

A "heterologous sequence" is one that originates from a foreign species, or, if from the same species, is substantially modified from its original form. For example, a promoter operably linked to a heterologous structural gene is from a species different from that from which the structural gene was derived, or, if from the same species, one or both are substantially modified from their original form.

A "glutenin polypeptide" is a gene product of a glutenin gene or glutenin polynucleotide sequence. A glutenin polypeptide can be either a LMW glutenin or a HMW glutenin. A glutenin polypeptide contains cysteine residues by which disulfide bonds are formed with other glutenin p lyp ptid s to form polymers. The composition and size of the repeat r gion is also important to polymer formation.

A "chimeric glutenin p lypeptide" is glutenin gene product that comprises a modified amino acid sequence. Modifications, as explained in detail b low, can be in the form of substitutions, deletions, or additions of singl amino acids or groups of amino acids. Thus, chimeric glutenin polypeptid s can be hybrid glutenin polypeptides comprising sequences from two or more different subunits or may be polypeptides in which single amino acid modifications are made.

In the case where an inserted polynucleotide sequence is transcribed and translated to produce a functional glutenin polypeptide, one of skill will recognize that because of codon degeneracy, a number of polynucleotide sequences will encode the same polypeptide. These variants are specifically covered by the terms "glutenin gene" or "qlutenin polynucleotide In addition, the terms specifically include thos full length sequences substantially identical (determined as described below) with a glutenin gene sequence and that encode proteins that retain the function of the glutenin polypeptide. Thus, in the case of wheat glutenin genes disclosed here, the term includes variant polynucleotide sequences which have substantial identity with the sequences disclosed here and which encode glutenin polypeptides capable of crosslinking by disulfide bonds with other glutenin polypeptides to form glutenin polymers.

Two polynucleotides or polypeptides are said to be "identical" if the sequence of nucleotides or amino acid residues, respectively, in the two sequences is the same when aligned for maximum correspondence as described below. The term "complementary to" is used herein to mean that the complementary sequence is identical to all or a portion of a reference polynucleotide sequence.

Sequence comparisons between two (or more) polynucleotides or polypeptides are typically performed by comparing sequences of the two sequences over a segment or "comparison window" to identify and compare local regions of sequence similarity. Optimal alignment of sequences for comparison may be conducted by the local homology algorithm of Smith and Waterman Adv. Appl. Math. 2: 482 (1981), by th homology alignment algorithm of Needleman and Wunsch J. Mol. Biol. 48:443 (1970), by the search for similarity method f Pearson and Lipman Proc. Natl.

(U.S.A.) Sci. 85: 2444 (1988), computerized Acad. by implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisc nsin Genetics Softwar Package, Genetics Computer Group (GCG), 575 Science Dr., Madison, WI), or by "Percentage inspection. of sequence identity" determined by comparing two optimally aligned sequences over a comparison window, wherein the portion of the polynucleotide sequence in the comparison window may comprise additions or deletions (i.e., gaps) as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical nucleic acid base or amino acid residue occurs in sequences to yield the number of matched positions, dividing number of matched positions by the total number positions in the window of comparison and multiplying the result by 100 to yield the percentage of sequence identity.

"substantial identity" polynucleotid of sequences means that a polynucleotide comprises a sequence that has at least 60% sequence identity, preferably at least 80%, more preferably at least 90% and most preferably at least 95%, compared to a reference sequence using the programs described above using standard parameters. One of skill will recogniz that these values can be appropriately adjusted to determine corresponding identity of proteins encoded by two nucleotide sequences by taking into account codon degeneracy, amino acid similarity, reading frame positioning and the Substantial identity of amino acid sequences for these purposes normally means sequence identity of at least 40%, preferably at least 60%, more preferably at least 90%, and most preferably at Polypeptides which are "substantially similar" share sequences as noted above except that residue positions which are not identical may differ by conservative amino acid Conservative amino acid substitutions refer to the interchangeability of residues having similar side chains. example, a group of amino acids having aliphatic side chains is glycine, alanine, valine, leucine, and isoleucine; a group of amino acids having aliphatic-hydroxyl side chains is serin and thr onine; a gr up of amino acids having amide-containing side chains is asparagin and glutamine; a group of amino acids

having aromatic side chains is phenylalanine, tyrosine, and tryptophan; a group of amino acids having basic side chains is lysine, arginine, and histidine; and a group of amino acids having sulfur-containing side chains is cystein and methionine. Pref rred conservative amino acids substitution groups are: valine-leucine-isoleucine, phenylalanine-tyrosine, lysine-arginine, alanine-valine, and asparagine-glutamine.

indication that nucleotide Another sequences ar substantially identical is if two molecules hybridize to each other under stringent conditions. Stringent conditions ar and will different sequence dependent be in different circumstances. Generally, stringent conditions are selected t be about 5°C to about 20°C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength and The Tm is the temperature (under defined ionic strength and pH) at which 50% of the target sequence hybridizes to a perfectly matched probe. However, nucleic acids which do not hybridize to each other under stringent conditions are still substantially identical if the polypeptides which they encode are substantially identical. This may occur, e.g., when a c py of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a diagram showing the domains of y-type and x-type HMW glutenins. The C- and N-terminal domains are shad d, the locations of cysteine residues are designated by S.

Figure 2A is a diagram showing the structure of a hybrid glutenin polypeptide of the invention.

Figure 2B is a diagram of the recombinant plasmid of th invention which encodes the polypeptide shown in Figure 2A.

Figure 3 is photograph of a gel stained with Coomassie blue showing the relative mobility of a hybrid HMW glutenin polypeptide of the invention (marked by the arrow).

DESCRIPTION OF THE PREFERRED EMBODIMENTS

This invention relates to plant glutenin genes, in particular wheat glutenin g n s. Nucleic acid sequ nces fr m glutenin genes can b us d to modify glut nin c ntent in transgenic plants, in particular wheat plants. In addition,

the inv ntion pr vides new se d-specific promoters useful for directing expression of desired genes in a number of plants.

Generally, the nomenclature and the laboratory pr cedures in recombinant DNA technology described b low ar those well known and commonly employed in the art. Standard techniques are used for cloning, DNA and RNA isolation, amplification and purification. Generally enzymatic reactions involving DNA ligase, DNA polymerase, restriction endonucleases and the like are performed according to the manufacturer's specifications. These techniques and various other techniques are generally performed according to Sambrook et al., Molecular Cloning - A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, (1989).

Glutenin polypeptides and genes

The HMW glutenins are composed primarily of a central domain of 45-90 repeating two or three simple motifs, comprising chiefly glutamine and proline. This unusual primary structure results in a rod-like secondary structure as assessed by circular dichroism and SEM. Cysteines in the C- and N-terminal domains of the molecules are known to be critical for the formation of intermolecular disulfide bonds (Figure 1). The quaternary interactions of these proteins are of interest because the disulfide crosslinks among HMW and LMW subunits is thought to be integral to the elastic properties of dough.

Any isolated glutenin gene can be used in the present invention. The particular polynucleotide sequence used is not a critical feature of the invention, so long as the desir d alteration in glutenin content is achieved. As noted above, HMW glutenins (both x-types and y-types) are encoded at th Glu-1 locus. In hexaploid wheat the group 1 chromosomes ar designated 1A, 1B and 1D. Studies of the frequency of allel s at each of the loci indicate the presence of at least three alleles at Glu-Al, 11 alleles at Glu-Bl, and six alleles at Glu-Dl. The gene products of each allele are designated by noting the chromosome, followed by its classification as x or y, and a number (see, MacRitchi, Advances in Food and Nutrition R search 36:1-87 (1992)).

Wheat HMW glutenin genes have b en cloned and described in the literature. For instance, a compl te s t of HMW glutenin

genes have ben isolated and s quenced fr m the hard red winter wheat Cheyenne (And rson et al. Wheat Genetics Symposium Proc. pp 699-704 (Bath Press, Cambridge, UK, 1988), including two, Dx5 and Dy10, with the high st correlation with good flour quality (Anderson et al. Nuc. Acids. Res. 17:461-462 (1989)). Other particularly useful HMW glutenins include Ax and Bx alleles.

The isolation of other glutenin genes may be accomplished by a number of techniques. For instance, oligonucleotide probes based on the sequences disclosed in the prior art can be used to isolate the desired gene from a cDNA or genomic DNA library. To construct genomic libraries, large segments of genomic DNA are generated by random fragmentation, e.g. using restriction endonucleases, and are ligated with vector DNA to form concatemers that can be packaged into the appropriate vector. To prepare a cDNA library, mRNA is isolated from endosperm and a cDNA library which contains the glutenin gene transcript is prepared from the mRNA.

Alternatively, the nucleic acids of interest can amplified from nucleic acid samples using amplification techniques. For instance, polymerase chain reaction (PCR) technology to amplify the sequences of the glutenin and related genes directly from genomic DNA, from cDNA, from genomic libraries or CDNA libraries. PCR and other amplification methods may also be useful, for example, to clon nucleic acid sequences that code for proteins to be expressed, to make nucleic acids to use as probes for detecting the presence of the desired mRNA in samples, for nucleic acid sequencing, or for other purposes. For a general overview of PCR see PCR Protocols: A Guide to Methods and Applications. (Innis, M, Gelfand, D., Sninsky, J. and White, T., eds., Academic Press, San Diego (1990)).

Polynucleotides may also be synthesized by well-known techniques as described in the technical literature. Se, e.g., Carruthers et al., Cold Spring Harbor Symp. Quant. Biol. 47:411-418 (1982), and Adams et al., J. Am. Chem. Soc. 105:661 (1983). Double stranded DNA fragments may then be obtained ither by synthesizing the complementary strand and annaling the strands tog ther under appr priate conditions, or by adding

the complementary strand using DNA polym ras with an appropriate primer sequence.

Isolated sequences pr par d as d scribed herein can then be us d to modify glutenin gen expression and theref re glutenin content in plants. One of skill will recognize that the nucleic acid encoding a functional glutenin protein need not have a sequence identical to the exemplified gen s disclosed here. Thus, genes encoding chimeric glut nin polypeptides can be used in the present invention.

above, glutenin polypeptides, like other noted proteins, have different domains which perform different Thus, the glutenin gene sequences need not be full functions. length, so long as the desired functional domains of th protein is expressed. Chimeric glutenin polypeptides can b readily designed utilizing various recombinant DNA techniques well known to those skilled in the art. For example, th chains can vary from the naturally occurring sequence at the primary structure level by amino acid substitutions, additions, deletions, and the like. In particular, cysteine residues may be added, deleted or moved within the polypeptide to achieve a modified glutenin polypeptide with desired properties. Chimeric polypeptides may also be produced by fusing coding sequences from two or more glutenin genes. All of these modifications can be used in a number of combinations to produce the final modified protein chain.

Preparation of recombinant constructs

To use isolated glutenin sequences in modifying glutenin content in plants, recombinant DNA vectors suitable transformation of plant cells are prepared. A DNA segu nce coding for the desired glutenin polypeptide, for example a cDNA or a genomic sequence encoding a full length protein, is conveniently used to construct a recombinant cassette which can be introduced into the desired plant. expression cassette will typically comprise the polynucleotide sequence operably linked to a promoter sequ no and other transcriptional and translational initiation regulatory s quences which will direct the transcription of the s quence from th glutenin g ne in the intended tissues (.g., nd sperm) of th transf rmed plant.

For exampl, a constitutive plant promoter fragment may be employed which will direct expression of the glutenin in all tissues of a plant. Such promoters are active under most nvironm ntal conditions and states of development or cell Examples of constitutive promoters include differentiation. mosaic cauliflower virus (CaMV) 35S transcription initiation region, the 1'- or 2'- promoter derived from T-DNA Agrobacterium tumafaciens, and other transcription initiation regions from various plant genes known to those of skill.

Alternatively, the plant promoter may be und r environmental control. Such promoters are referred to here as "inducible" promoters. Examples of environmental conditions that may effect transcription by inducible promoters includ pathogen attack, anaerobic conditions, or the presence of light.

Typically, the promoters used in the constructs of the invention will be "tissue-specific" and are under developmental control such that the desired gene is expressed only in certain tissues, such as leaves, roots, fruit, seeds, or flowers. Promoters that direct expression in seeds, particularly the endosperm are particularly preferred. Examples of such promoters include the promoter from genes encoding seed storage proteins, such as napin, cruciferin, phaseolin, and the like (see, U.S. Patent No. 5,420,034). Other promoters suitable for expressing glutenin genes in cereals include promoters from genes encoding gliadins, cereal prolamines (e.g., zein, hordein, secalin, and avenin) and starch biosynthetic enzymes.

endogenous promoters from glutenin genes particularly useful for directing expression of glutenin gen s to the seed, particularly the endosperm. These seed-specific promoters can also be used to direct expression of heterologous structural genes. Thus, the promoters can be used recombinant expression cassettes to drive expression of any gene whose expression in seeds is desirable. Examples include genes encoding proteins useful in increasing the nutritional value of seeds (e.g., genes ncoding proteins involved in lipid, protein, and carbohydrate or starch biosynthesis). Oth r g n s includ those ncoding pharmaceutically us ful

compounds, and genes encoding plant resistance products to combat fungal or oth r infections of the seed.

glut nin promoters can also be used to initiat transcription of mRNA molecules to inhibit expression of an endogenous endosperm gene. Exemplary genes whose expression could be inhibited include genes encoding glutenins, enzym s involved in carbohydrate or lipid synthesis, secalins, and the Means for inhibiting gene expression in plants using recombinant DNA techniques are well known. For instance, antisense technology can be conveniently used. see, e.g., Sheehy et al., Proc. Nat. Acad. Sci. USA, 85:8805-8809 (1988), and Hiatt et al., U.S. Patent No. 4,801,340. Catalytic RNA molecules or ribozymes can also be used to inhibit expression of endosperm-specific genes. The design and use of target RNAspecific ribozymes is described in Haseloff et al. Nature, 334:585-591 (1988). Introduction of nucleic acid configured in the sense orientation has also been shown to be an effective means by which to block the transcription of target genes. an example of the use of sense suppression to modulate expression of endogenous genes see, Napoli et al., The Plant 2:279-289 (1990), and U.S. Patents Nos. 5,034,323, 5,231,020, and 5,283,184.

The glutenin promoters of the invention are typically at least about 400 base pairs in length, and often at least about 800 or about 1000 base pairs. The length of the promoters is typically less than about 3500 base pairs, usually less than about 2800 base pairs and often less than about 2000 base pairs in length. The length of the promoters is counted upstream from the translation start codon of the native gene. One of skill will recognize that use of the "about" to refer to lengths of nucleic acid fragments is meant to include fragments of various lengths that do not vary significantly from the lengths recited here and still maintain the functions of the claimed promoters (i.e., seed-specific gene expression).

To identify glutenin promoters, the 5' portions of a genomic glutenin gene clone is analyzed for sequences characteristic of promoter s quences. For instance, promoter sequence elements includ the TATA box cons nsus sequence (TATAAT), which is usually 20 t 30 bas pairs upstream of the transcription start site. In plants, further upstream fr m the

TATA box, at positions -80 to -100, there is typically a promoter element with a series of adenines surrounding the trinucleotid G (or T) N G. J. Messing et al., in Genetic Engineering in Plants, pp. 221-227 (Kosage, Meredith and Hollaender, eds. 1983).

In preparing expression vectors of the invention, sequences other than the promoter and the structural gene of interest are also preferably used. If proper polypeptid expression is desired, a polyadenylation region at the 3'-end of the glutenin coding region should be included. The polyadenylation region can be derived from the natural gen, from a variety of other plant genes, or from T-DNA.

The vector comprising the sequences from a glutenin gene will typically comprise a marker gene which confers a selectable phenotype on plant cells. For example, the marker may encode biocide resistance, particularly antibiotic resistance, such as resistance to kanamycin, G418, bleomycin, hygromycin, or herbicide resistance, such as resistance to chlorosluforon, or phosphinothricin (the active ingredient in bialaphos and Basta).

Preparation of transgenic plants

The DNA constructs described above may be introduced into the genome of the desired plant host by a variety of conventional techniques. Techniques for transforming a wide variety of higher plant species are well known and described in the technical and scientific literature. See, for example, Weising et al., Ann. Rev. Genet. 22:421-477 (1988).

The DNA construct may be introduced directly into the genomic DNA of the plant cell using techniques such as biollistic methods, electroporation, PEG poration, and microinjection of plant cell protoplasts or embryogenic callus. Alternatively, the DNA constructs may be combined with suitable T-DNA flanking regions and introduced using an Agrobacterium tumefaciens or A. rhizogenes vector.

Particle bombardment techniques are described in Klein t al., Nature 327:70-73 (1987). A particularly preferred m thod of transforming wh at and other cereals is the bombardment f calli deriv d from immature embryos as d scrib d by W ks et al. Plant Physiol. 102:1077-1084 (1993).

Microinjection techniques are known in the art and well described in the scientific and patent literature. The introduction of DNA constructs using polyethylene glycol precipitation is described in Paszkowski et al. Embo J. 3:2717-2722 (1984). Electroporation techniques are described in Fromm et al. Proc. Natl. Acad. Sci. USA 82:5824 (1985).

Agrobacterium tumefaciens-meditated transformation techniques also are well described the in scientific literature. See, for example Horsch et al. Science 233:496-498 (1984), and Fraley et al. Proc. Natl. Acad. Sci. USA 80:4803 (1983). Although Agrobacterium is useful primarily in dicots, certain monocots can be transformed by Agrobacterium. instance, Agrobacterium transformation of rice is described by Hiei et al. Plant J. 6:271-282 (1994).

The present invention is particularly useful in wheat and A number of methods of transforming cereals other cereals. been described in the literature. For instance, transformation of rice is described by Toriyama Bio/Technology 6:1072-1074 (1988), Zhang et al. Theor. Appl. Gen. 76:835-840 (1988), and Shimamoto et al. Nature 338:274-276 Transgenic maize regenerants have been described by Fromm et al., Bio/Technology 8:833-839 (1990) and Gordon-Kamm et al., Plant Cell 2:603-618 (1990)). Similarly, oats (Somm rs et al., Bio/Technology 10:1589-1594 (1992)), wheat (Vasil et al., Bio/Technology 10:667-674 (1992)); Weeks et al., Plant Physiol. 102:1077-1084 (1993)), sorghum (Casas et al., Proc. Natl. Acad. Sci. USA 90:11212-11216 (1993)), rice (Li et al., Plant Cell Rep. 12:250-255 (1993)), barley (Yuechun and Lemaux, Plant Physiol. 104:37-48 (1994)), and rye (Castillo et al., Bio/Technology 12:1366-1371 (1994)) have been transformed via bombardment.

Transformed plant cells which are derived by any of the above transformation techniques can be cultured to regenerate a whole plant which possesses the transformed genotype and thus the desired phenotype. Such regeneration techniques rely on manipulation of certain phytohormones in a tissue culture growth medium, typically relying on a biocide and/or herbicid mark r which has be n introduced together with th glutenin polynucleotide s quences. Plant r g neration fr m cultured protoplasts is described in Evans et al., Protoplasts Isolation

and Culture, Handbook of Plant Cell Culture, pp. 124-176, MacMillilan Publishing Company, New York, 1983; and Binding, Regeneration of Plants, Plant Protoplasts, pp. 21-73, CRC Press, Boca Rat n, 1985. Regeneration can also be obtained from plant callus, explants, organs, or parts thereof. Such regeneration techniques are described generally in Klee et al. Ann. Rev. of Plant Phys. 38:467-486 (1987).

The methods of the present invention are particularly useful for incorporating the glutenin polynucleotides into transformed plants in ways and under circumstances which are not found naturally. In particular, the glutenin polypeptid s may be expressed at times or in quantities which are not characteristic of natural plants.

One of skill will recognize that after the expression cassette is stably incorporated in transgenic plants and confirmed to be operable, it can be introduced into other plants by sexual crossing. Any of a number of standard breeding techniques can be used, depending upon the species to be crossed.

The invention has use over a broad range of types of plants. The glutenin genes are preferably expressed in cereal species commonly used for production of flour, e.g., wheat, rye, oats, and the like.

The seed-specific promoters from glutenin genes can be used in essentially any plant species. For instance, the promoters or genes described here can be used in species from genera Asparagus, Atropa, Avena, Brassica, Citrullus, Capsicum, Cucumis, Cucurbita, Daucus, Fragaria, Glycine, Gossypium, Helianthus, Heterocallis, Hyoscyamus, Juglans, Lactuca, Linum, Lolium, Lycopersicon, Malus, Manihot, Majorana, Medicago, Nicotiana, Oryza, Panieum, Pannesetum, Persea, Phaseolus, Pisum, Pyrus, Prunus, Raphanus, Secale. Senecio, Sinapis, Solanum, Sorghum, Triticum, Vitis, Vigna, and, Zea. Plants that are particularly useful in the invention include Avena, Brassica, Glycine, Hordeum, Oryza, Phaseolus, Pisum, Secale, Sorghum, Triticum, Vigna, and Zea.

As noted above, the invention is particularly useful for improvement of wheat cultivars. Any wheat cultivar may be improved by the methods of the invention. Exemplary lines

include hexapl id lines such as Anza, Shasta, Yecoro Rojo, Siouxland, Fr dom, Tam107, and Tam200.

The ffect of the modification of glutenin gene xpressi n can b measured by detecti n of increases or decr as s in desired glutenin protein levels using, for instance, gel electrophoresis, as described below. Quantification of HMW glutenin content can be carried out by SDS/PAGE densitometry, as described below.

Other methods for quantifying glutenin content include sonication of flour dispersions in SDS buffer to solubilize glutenin and other relatively insoluble proteins (see, Singh and MacRitchie in Wheat End-Use Properties: Wheat and Flour Characterization for Specific End-Uses H. Salovaara, ed., pp 321-326 (University of Helsinki, Helsinki, 1989). This method is useful in solubilizing at least 95% of total flour protein. Quantification of the glutenin fraction can then be performed using size exclusion high performance liquid chromatography as described by Batey et al., Cereal Chem. 68:207-209 (1991). Other suitable methods include reverse phase HPLC and capillary electrophoresis.

The plants of the invention have increased in glutenin content in the endosperm of mature seeds as compared to the parental lines from which they are derived. The increase is usually at least about 15% as compared to the parental lin. More usually, the increase is at least about 20%, preferably at least about 35%, and in more preferred embodiments at least about 45%. Any of the methods for quantifying glutenins in the endosperm described above can be used to determine the percent increase in glutenin content. A wheat seed typically reaches maturity about 6 to about 10 weeks after anthesis.

Another measure of increased glutenin content in transgenic plants of the invention is glutenin content as a percent of total protein of a mature seed or flour derived from the mature seed. Different wheat cultivars have different glutenin contents as measured using standard methods. Thus, the final glutenin content of the plants of the invention will depend upon the parental line. In the case where HMW glutenin genes are used, the plants f th invention will typically have HMW glutenin contents of at least about 15%, usually at least

about 18%, and pr ferably at 1 ast about 20% as measur d using standard methods such as those described in detail below.

Analysis of the rheological properties of flour derived from the transgenic plants of the inventi n can be carried out according to standard physical dough-testing instruments widely used to measure flour and dough quality (see, e.g., MacRitchie, Advances in Food and Nutrition Research 36:1-87 (1992)). methods include, for instance, use of extensographs to measure tensile strength. Two of the main parameters measured are maximum resistance (R_mx) and extensibility (Ext). include mixographs and bake-test loaf (MacRitchie and Gras Cereal Chem. 50:292-302 (1973) as well as SDS-sedimentation tests, amelographs, and cookie methods.

The following Examples are offered by way of illustration, not limitation.

Example 1

This example demonstrates increase in glutenin content in transgenic wheat plants comprising recombinant constructs of the invention.

MATERIALS AND METHODS

Construction of the hybrid HMW glutenin expression plasmid

Two HMW glutenin subunits, Dx5 and Dy10, encoded by the Glu-D1-1b and Glu-D1-2b genes, respectively, have been most often associated with superior dough strength by genetic correlation studies. A bacterial expression vector comprising a fusion of coding regions of the Dy10 and Dx5 genes is described by Shani et al., Plant Physiol. 98:433-441 (1992). The hybrid HMW glutenin polypeptide encoded by this gene is diagrammed in Figure 2A.

In this example, the Dy10:Dx5 hybrid coding region was reunited with 5' and 3' flanking sequences from the native Dy10 and Dx5 genes. The construction of a fusion at the HindIII site located at base pairs 429-435 and 385-391 after the A of the start codons of the Glu-D1-2b (Genbank Accession N. X12929) and Glu-D1-1b (G nbank Accession No. X12928) genes (And rson et al. Nuc. Acids Rs. 17:461-462 (1989)) and its

insertion into the bact rial expr ssion vector pet3A to form pet-3a-10/5 is describ d by Shani et al., supra.

native Dyl0 g ne provides the the transcription start site, the 5' transcribed untranslat d region and the first 145 codons including the 21 amino acid signal peptide. The native Dx5 gene provides its codons 130 to 848 followed by two translation stop codons, the transcripti n termination and poly(A) addition signals. Because amino acids 138 through 147 are shared by the Dy10 and Dx5 glutenin subunits, the central region of repeating amino acid motifs of the hybrid glutenin polypeptide (thickest boxes in Figure 2A) identical to that of Dx5. The configuration of th cysteines (S) and thus of potential disulfide bonds is chimeric in nature: the N-terminal cysteines are of the y-type and th single C-terminal cysteine is typical of x-type subunits.

order to reconstitute the promoter transcription start and termination sites, and any other regulatory signals needed for expression in wheat, fragments were isolated and combined in a single ligation reaction with the Bluescript KS (Stratagene) vector cut with EcoRI and BamHI: 1) the 715 bp StuI/DrdI fragment that contains the junction region in pET-3a-10/5; 2) the 2800 bp fragment from a clone of the native Dy10 gene beginning at the EcoRI site 5' to the gene and ending at the StuI site 74 bp after the A in the start codon; and 3) the 3800 bp fragm nt from a clone of the native Dx5 gene that starts at the DrdI site 744 bp after the A in the start codon and ends at a BamHI site in the vector just outside the EcoRI site in the 3' flanking region. A plasmid with the correct structure was identified by restriction analysis and named pGlu10H5 (Figur This plasmid was deposited under the terms of the Budapest Treaty in the Agricultural Research Culture Collection (NRRL), 1815 North University Street, Peoria, Illinois, USA, on January 12, 1996 and has been assigned Accession No. NRRL B-21517.

The plasmid UBI:BAR which comprises the maize ubiquitin promoter operably linked to the bar g ne encoding the nzym PAT, which inactivats phosphinothricin was prepared as described Corn jo et al. Plant Mol. Biol. 23:567-581 (1993).

Both plasmids wer prepard by the alkalin lysis method using a Qiagen kit for the final purificati n. The DNAs were stored in TE buffer at a concentration of 1 mg/ml.

Wheat Transformation

12.5 ug of each DNA (an approximately 2:1 molar ratio of UBI:BAR to pGlu10H5) was coated onto gold particles for bombardment into immature wheat embryos of cultivar Bobwhite essentially as described by Weeks et al. Plant Physiol. 102:1077-1084 (1993) except that the embryos were incubated on callus induction media containing mannitol for 4 hours before and 20 hours after bombardment. Transformants were selected on the basis of their resistance to lmg/L bialaphos at the callus and green shoot stages of regeneration and to 3 mg/L bialaphos at the rooting stage of regeneration as described by Weeks t al., supra.

Protein Analysis

Protein extracts from immature wheat endosperm wer prepared by grinding the tissue or cells in SDS-PAGE sample loading buffer [66 mM Tris pH 6.8, 3% (w/v) SDS, 0.05% (w/v) Bromphenol blue, 2% (v/v) b-mercaptoethanol]. Protein extracts from mature dry wheat seeds were prepared by dissolving flour sample buffer. SDS-PAGE Was according to standard in procedures. The separation gels for detection and quantitation of the hybrid subunit were 10% (w/v) acrylamide 0.05% (w/v)bis-acrylamide. The gels were run at 40 mamps constant current until 20 minutes after the bromphenol blue dye front had run off the gel (about 1000 Volt-hours).

The gels were stained by the Neuhoff method with a colloidal suspension of Coomassie blue. Protein standards for molecular weight calibration were purchased from Pharmacia and consisted of Phosphorylase b (94 kDa) Bovine Serum Albumin (67 kDa), Ovalbumin (43 kDa), Carbonic Anhydrase (30 kDa), Soybean Trypsin Inhibitor (20.1 kDa) and a-Lactalbumin (14.4 kDa). Stained gels were scanned using an Alpha Innotech densitometer.

DNA Analysis

Southern analyses of g nomic DNA was as describ d in Weeks et al., supra, except that 10 ug of DNA wer loaded into each

slot of a 0.6% agaros g 1. The probe consisted of the 1578 bp PvuII fragment from the coding region of Dx5 (bracketed in Figure 2B).

RESULTS

Forty-five independent callus pieces were selected from 4980 embryos bombarded in 10 different experiments; each yielded one to five plants. The To primary regenerated plants and the lines derived from them are designated with an unique number following the bombardment number, eq., plant 126 regenerated from bombardment 7 is 7-126. Plants from twenty-six of these independent line gave rise to progeny embryos that were able to germinate in the presence of 3 mg/L bialaphos when excised from immature seeds three to four weeks after anthesis.

The immature endosperm tissue corresponding to each excised embryo was extracted and screened by SDS-PAGE for th presence of the hybrid glutenin. This SDS-PAGE analysis showed that the migration of the hybrid HMW glutenin subunit is clearly distinguishable from the native HMW subunits under these gel conditions.

Although the lanes did not contain the same amounts of total protein, it was evident that there is variation in the expression levels of the hybrid subunit relative to those of the native HMW subunits in the same seed. Of the twenty-six resistant lines that were derived from co-bombardm nt experiments including pGlu10H5, sixteen (60%) lines showed expression of the hybrid subunit in the T₁ generation.

Expression stability

At each successive generation, 24 or 25 immature embryos 3-4 weeks after anthesis were tested for their ability to germinate on bialaphos. Plants were classified as homozygous if all 25 of their progeny embryos exhibited resistance. levels of transgene expression were assessed in successiv generations. In most cases, expression was stable. An apparent increase expression between the in T1 T2 generations in some plants was explain d by the incr as gene dosage between the heterozyg us and homozygous prog ny of the TO plant and T1 g nerati n, respectively. the end sperm is triplicate, individual selfed pr geny of the T_{o}

h terozygote could have 0, 1, 2 or 3 copi s and th T1 lane contains an average of 8 of these individuals.

In four of the 15 wh at lines analyzed, expr ssion of the hybrid HMW glutenin has not remained at the same 1 v ls, but rather has declined as homozygous resistant lines were selected in the T_2 or the T_3 generations. The decline could be du either to hybrid HMW glutenin transgene inactivation or to loss of unlinked pGlulOH5 transgene(s) copies via segregation.

Cross-linking ability of the hybrid subunit

As noted above, the hybrid HMW glutenin contains a uniqu configuration of cysteines, y-type at the N-terminus and x-type at the C-terminus. The hybrid HMW glutenin behaved exactly like the native subunits in terms of its solubility in SDS buffers with and without a reducing agent, when examined by SDS-PAGE. The native and hybrid HMW glutenins were partially extracted from both the Bobwhite and the transgenic flours by SDS alone and were present in lower molecular weight polymers. The majority of the hybrid glutenin, however, was insoluble in SDS buffer and could only be extracted when a reducing agent Thus, the configuration of cysteines in the hybrid glutenin does not preclude the formation of either intermolecular disulfide bonds or high molecular weight polymers.

Relative expression levels

Figure 3 shows SDS-PAGE analysis of protein extracts from eight different transgenic lines (Lanes 1-8) and Bobwhite (C). Each sample was prepared from flour of a mixture of eight mature seeds of a single homozygous T, or T, plant. Under thes gel conditions, the native By9 and Dy10 subunits are not well resolved and run as a closely spaced doublet of apparent molecular weight 94 kDa. Likewise the two isoforms of the Dx5 subunit are not well-resolved and migrate as a broad band. The Bx7 band is well-separated from other proteins and the darkest of the HMW subunits in intensity. In all the lines except th one shown in Lane 3, the hybrid HMW subunit is present in at least as high a quantity as the native Ax2* (largest) HMW subunit. Thus wen a single copy of the hybrid transg ne (Lane

1) can support significant expression 1 vels, comparable to those of th native HMW subunits.

In on of the lines (shown in lane 2), synthesis of native HMW glutenin synthesis has declined r lativ to the other proteins. The extract shown there exhibits reduced synthesis of all the native HMW glutenin compared to the lower molecular weight seed proteins in the same lane.

In order to estimate the levels of the hybrid HMW glutenin relative to those of the native HMW glutenin and other seed proteins, the gel was scanned by densitometry. All the wh at prolamines stay on the gel. However, since proteins f molecular weights less than 33 kDa are run off the gel und r the conditions used to separate the hybrid HMW glutenin from the nearby Dx5 subunit band, only relative quantities could be assessed. The results of these comparisons are shown in Table I along with the number of intact transgene copies estimated from restriction enzyme digests of genomic DNA (see below). The hybrid HMW glutenin band constitutes more than 20% of the total HMW glutenins in all the lines except that in lane 3. Thus the transgene has made a significant contribution to the HMW glutenin composition in all the transgenic lines.

In the case where endogenous HMW glutenin accumulation is reduced (Lane 2), the hybrid makes up 62% of the total complement of HMW glutenin. In most lines, the sum of the HMW glutenin accumulation is elevated 29 to 45% relative to the HMW glutenin content of Bobwhite (all sums are normalized to the remainder of the protein their respective lanes). Thus this quantitative analysis suggests that addition of HMW-glutenin gene copies raises the levels of HMW glutenin accumulation in mature seeds.

DNA analysis

DNA was isolated from the leaves of transgenic plants of each line and assayed for the presence of transgenes by genomic Southern analysis, as described above. Hybridizing bands known to correspond to the transgenes were scanned and thir densities compared. The results are shown in Table I. The number ranges from 1 in line 11-291 to about 6 in 14-23 (Table I). This range is much broader than that in total HMW glut nin levels or even hybrid HMW glutenin 1 vels. How v r the most

highly xpressing lin had one of th highest copy numbers (Line 4) and th poor st expression was in a single copy line (Line 3).

Table I. Transgenic HMW-glutenin gene activity in individual wheat lines.

Line	Transgene Expression	Total HM subunit as % of total	
1	23%	135%	1
2	62%	70%	5-6
3	15%	145%	1
4	42%	145%	4-5
5	23%	140%	2
6	24%	129%	2
7	22%	135%	3-4
8	20%	143%	3-4

DISCUSSION

The experiments reported here show that levels of HMW glutenins can be increased by addition of gene copies. The behavior of the hybrid subunit is indistinguishable from that of the native subunits. The protein accumulates in the endosperm tissue over the course of seed development and is cross-linked by disulfide bonds into large polymers that cannot be extracted by SDS without addition of reducing agents.

Expression of the hybrid glutenin is under the control f the native Dy10 5' and Dx5 3' flanking sequences. Thes sequences were sufficient to achieve high levels of the modified seed storage protein. Even a single gene copy supported expr ssion l vels comparable to thos of the native HMW glutenin genes. Thus, these sequences s rve as a source of effective regulatory sequences suitable for xpr ssion of other

proteins in transgenic wheat and potentially other cereal ndosperm.

Exampl 2

This example describes the transformation of wheat using a recombinant construct encoding Dx5 and Dy10.

The recombinant expression vectors were prepared as generally as described above except that the complete Dx5 or Dy10 gene, including 5' and 3' regulatory sequences, were us d. The constructs were then introduced into immature wheat embryos by particle bombardment as described above. Analysis of T_0 plants and their progeny is performed as described above.

The above examples are provided to illustrate the invention but not to limit its scope. Other variants of the invention will be readily apparent to one of ordinary skill in the art and are encompassed by the appended claims. All publications, patents, and patent applications cited herein are hereby incorporated by reference for all purposes.

Applicant's or agent's file	International application No
reference number A700/363	

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indica	ations made below relate to the microorganism refere	ed to in the description						
on page	17 , line 31							
B. IDENTIF	FICATION OF DEPOSIT	Further deposits are identified on an additional sheet						
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	positary institution (including postal code and count University Street pis 61804	ליס						
Date of depos	sit	Accession Number						
January 12,	1996	NRRL B-21517						
C. ADDITI	ONAL INDICATIONS (leave blank if not applicable	This information is continued on an additional sheet						
a sample of European P	1 microorganism is disclosed in the specification. In respect to those designations in which a European Patent is sought a sample of the deposited microorganism will be made available until the publication or the mention of the grant of the European Patent or until the date on which the application has been refused or withdrawn or is deemed to be withdrawn, only the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC).							
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WHAT IS CLAIMED IS:

1. A m thod of alt ring glut nin content in the endosperm of a wh at plant, the method comprising:

stably transforming a parental wheat plant with a recombinant expression cassette comprising a nucleic acid encoding a glutenin polypeptide and a seed-specific promoter to produce a stably transformed wheat plant; and

selecting a progeny of said stably transformed wheat plant having a glutenin content in the endosperm of a mature seed at least about 30% different than the glutenin content in the endosperm of a mature seed from the parental wheat plant.

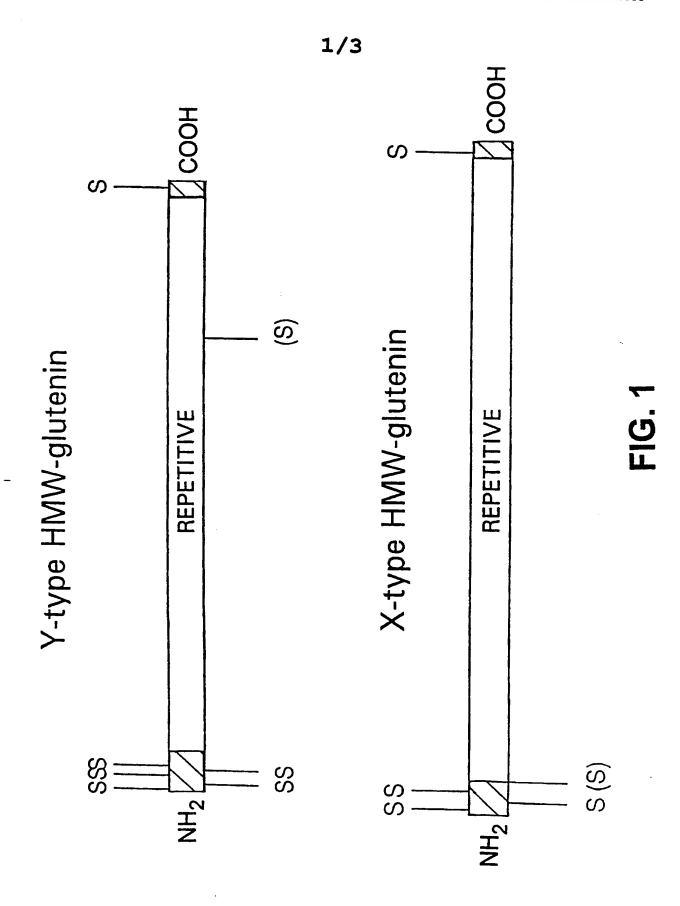
- 2. The method of claim 1, wherein the nucleic acid is stably transformed into the first wheat plant using particle bombardment.
- 3. The method of claim 1, wherein the nucleic acid encodes a chimeric glutenin polypeptide.
- 4. The method of claim 1, wherein the chimeric polypeptide comprises sequences from an x-type glutenin polypeptide and a y-type glutenin polypeptide.
- 5. The method of claim 1, wherein the x-type polypeptide is encoded by the Glu-Dl-lb gene.
- 6. The method of claim 1, wherein the y-type polypeptide is encoded by the Glu-D1-2b gene.
- 7. The method of claim 1, wherein the promoter is from the Glu-D1-2b gene.
- 8. The method of claim 1, wherein the parental wh at plant is Bobwhite.
- 9. A wheat plant comprising a recombinant expression cassett comprising a nucleic acid encoding a glut nin polypeptide and a seed-specific promoter, the plant having a

glutenin c ntent in the end sperm of a mature sed at last about 30% different than the glutenin content in the endosperm of a mature seed from a parental wheat plant.

- 10. The plant of claim 9, wherein the nucleic acid encodes a chimeric glutenin polypeptide.
- 11. The plant of claim 10, wherein the chimeric polypeptide comprises sequences from an x-type polypeptide and a y-type glutenin polypeptide.
- 12. The plant of claim 11, wherein the x-type polypeptid is encoded by the Glu-D1-1b gene.
- 13. The plant of claim 11, wherein the y-type polypeptid is encoded by the Glu-D1-2b gene.
- 14. The plant of claim 9, wherein the promoter is from the Glu-Dl-2b gene.
- 15. The plant of claim 9, wherein the high molecular weight (HMW) glutenin accounts for at least about 15% of the total protein in the endosperm of the mature seed.
- 16. The plant of claim 9, wherein the parental wheat plant is Bobwhite.
 - 17. A seed derived from the plant of claim 9.
- 18. A recombinant construct comprising a wheat glutenin gene promoter of about 400 to about 2800 nucleotides operably linked to a heterologous polynucleotide sequence encoding a chimeric glutenin polypeptide, wherein the chimeric glutenin polypeptide is encoded by polynucleotide sequences from the Glu-Dl-lb gene and the Glu-Dl-2b gene.
- 19. The recombinant construct of claim 18, wh rein the wheat glutenin gene is Glu-D1-2b.

20. A plant comprising th recombinant construct of claim 18.

21. Wheat flour derived from th plant of claim 1.



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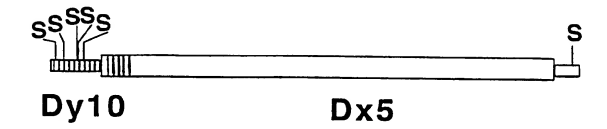


FIG. 2A

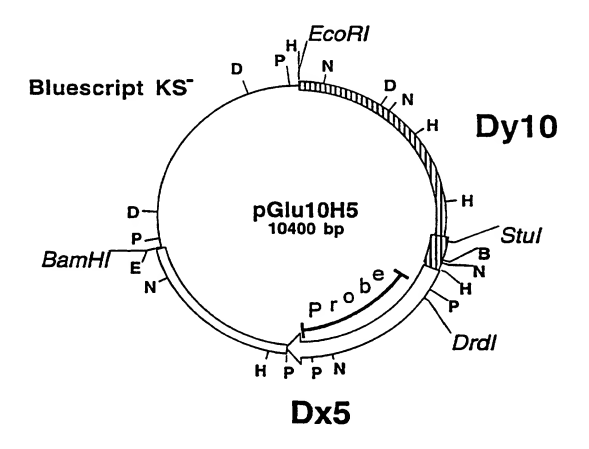


FIG. 2B

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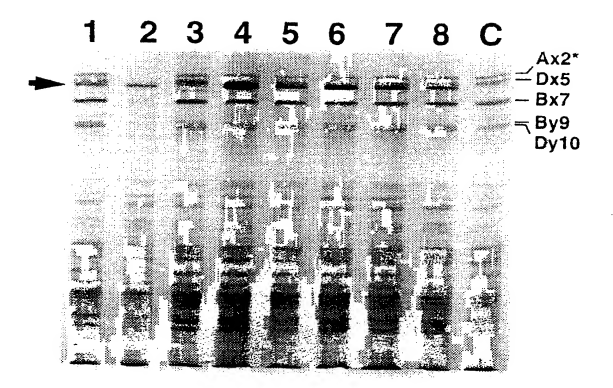


FIG. 3

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/00510

A. CLASSIFICATION OF SUBJECT MATTER						
IPC(6) :Please See Extra Sheet.						
According t	US CL: Please See Extra Sheet. According to International Patent Classification (IPC) or to both national classification and IPC					
8. FIEL	DS SEARCHED					
Minimum d	ocumentation searched (classification system follower	d by classification symbols)				
U. S . :	800/200, 205, 250, DIG 58; 435/172.3, 240.4, 320.	1; 536/23.6, 24.1				
Documentat	ion searched other than minimum documentation to the	e extent that such documents are included	in the fields searched			
Electronic d	lata base consulted during the international search (na	ame of data base and, where practicable.	search terms used)			
	BA, MEDLINE, CAPLUS, GENBANK, BIOSIS erms: wheat, glutenin, transformation, Glu-D1-	2b, Giu-D1-1b				
C. DOC	CUMENTS CONSIDERED TO BE RELEVANT					
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.			
Y	SHANI et al. Role of amino- and countries the folding and oligomerization weight glutenin subunits. Plant pages 433-441, see entire docum	of wheat high molecular Physiol. 1992, Vol. 98,	1-21			
Y	WEEKS et al. Rapid production of multiple independent lines of fertile transgenic wheat (<i>Triticum aestivum</i>). Plant Physiol. 1993, Vol. 102, pages 1077-1084, especially pages 1078-1079.					
Y	1-21					
X Furt	ner documents are listed in the continuation of Box C	See patent family annex.				
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/00510

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT						
Category*	Citation of document, with indication, where appropriate, of the relev	ant passages	Relevant to claim No			
Y	GALILI, G. Heterologous expression of a wheat high weight glutenin gene in <i>Escherichia coli</i> . Proc. Natl. USA. October 1989, Vol. 86, pages 7756-7760, see endocument.	Acad. Sci.	1-21			
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/00510

A. CLASSIFICATION OF SUBJECT MATTER: IPC (6):
C12N 15/00, 15/05, 15/09, 15/11, 15/29, 15/64, 15/82; A01H 1/00, 3/00, 4/00
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800/200, 205, 250, DIG 58; 435/172.3, 240.4, 320.1; 536/23.6, 24.1

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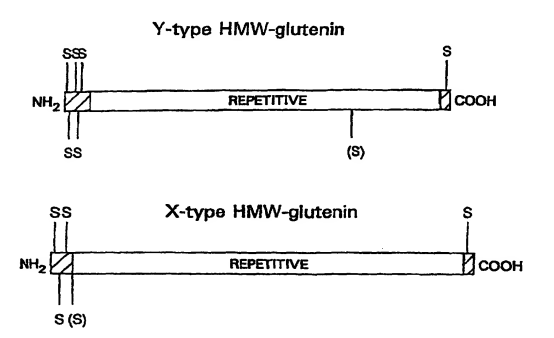
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(51) International Patent Classification 6:		(11) Internati nal Publication Number:	WO 97/25419		
C12N 15/00, 15/05, 15/09, 15/11, 15/29, 15/64, 15/82, A01H 1/00, 3/00, 4/00	A1	(43) International Publication Date:	17 July 1997 (17.07.97)		
 (21) International Application Number: PCT/US (22) International Filing Date: 10 January 1997 ((30) Priority Data: 08/586,331 16 January 1996 (16.01.96 (71) Applicant: THE UNITED STATES OF AMERICA sented by TH [US/US]; E SECRETARY OF ACTURE, Washington, DC 20250 (US). (72) Inventors: BLECHL, Ann, E.; 1005 Buchanan Street, CA 94706 (US). ANDERSON, Olin, D.; 208 Street, Pleasant Hill, CA 94523 (US). (74) Agents: PENDORF, Stephan, A. et al.; Suite 1000 Westshore Boulevard, Tampa, FL 33609 (US). 	10.01.9 A, repr GRICUI , Alban Posha	BY, CA, CH, CN, CZ, DE, DK IL, IS, JP, KE, KG, KP, KR, KZ LV, MD, MG, MK, MN, MW, RU, SD, SE, SG, SI, SK, TJ, T VN, ARIPO patent (KE, LS, M patent (AM, AZ, BY, KG, KZ, N patent (AT, BE, CH, DE, DK, I LU, MC, NL, PT, SE), OAPI p CM, GA, GN, ML, MR, NE, SI Published With international search report	I, EE, ES, FI, GB, GE, HU, Z, LC, LK, LR, LS, LT, LU, MX, NO, NZ, PL, PT, RO, TM, TR, TT, UA, UG, UZ, IW, SD, SZ, UG), Eurasian MD, RU, TJ, TM), European ES, FI, FR, GB, GR, IE, IT, Patent (BF, BJ, CF, CG, CI, N, TD, TG).		
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(54) Title: GLUTENIN GENES AND THEIR USES



(57) Abstract

The present invention is directed to methods of improving flour by altering glutenin content in seeds of plants, particularly wheat. The invention relates to methods of introducing a recombinant construct comprising a glutenin gene into a parental plant.

^{* (}Referred to in PCT Gazette No. 43/1997, Section II)

^{** (}Referred to in PCT Gazette No. 49/1997, Section II)

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